Calibrating the Cryogenian

Francis A. Macdonald,1,* Mark D. Schmitz,2 James L. Crowley,2 Charles F. Roots,3 David S. Jones,4 Adam C. Maloof,5 Justin V. Strauss,6 Phoebe A. Cohen,1 David T. Johnston,1 Daniel P. Schrag1

The Neoproterozoic was an era of great environmental and biological change, but a paucity of direct and precise age constraints on strata from this time has prevented the complete integration of these records. We present four high-precision U-Pb ages for Neoproterozoic rocks in northwestern Canada that constrain large perturbations in the carbon cycle, a major diversification and depletion in the microfossil record, and the onset of the Sturtian glaciation. A volcanic tuff interbedded with Sturtian glacial deposits, dated at 716.5 million years ago, is synchronous with the age of the Franklin large igneous province and paleomagnetic poles that pin Laurentia to an equatorial position. Ice was therefore grounded below sea level at very low paleolatitudes, which implies that the Sturtian glaciation was global in extent.

Middle Neoproterozoic or Cryogenian strata [850 to 635 million years ago (Ma)] contain evidence for the breakup of the supercontinent Rodinia, widespread glaciation (1, 2), high-amplitude fluctuations in the carbon cycle, a major diversification and depletion in the microfossil record, and the onset of the Sturtian glaciation. A volcanic tuff interbedded with Sturtian glacial deposits, dated at 716.5 million years ago, is synchronous with the age of the Franklin large igneous province and paleomagnetic poles that pin Laurentia to an equatorial position. Ice was therefore grounded below sea level at very low paleolatitudes, which implies that the Sturtian glaciation was global in extent.

We present four high-precision U-Pb isotope dilution–thermal ionization mass spectrometry (ID-TIMS) dates from intrusive and volcanic rocks within Neoproterozoic strata of northwestern Canada. These dates, coupled with high-resolution δ13C profiles (11), allow us to synthesize Cryogenian geological, geochemical, paleomagnetic, and paleontological data both regionally and globally. The accurate integration of these records places hard constraints on the timing and extent of the Sturtian glaciation and its relationship to the Franklin large igneous province (LIP) and the Cryogenian microfossil record.

Neoproterozoic strata are exposed in erosional windows (inliers) through Paleozoic carbonate rocks in northwestern Canada for more than 1500 km, from the Alaska border to the Atlantic Ocean (12). The snowball Earth hypothesis (1, 2) was developed in response to strong paleomagnetic evidence for low-latitude glaciation from the Elatina Formation in Australia (5, 6). The Elatina Formation and its distinct cap carbonate have been correlated with chemo- and lithostratigraphy to Marinoan-age glacial deposits in the Ghaub Formation in Namibia (635.5 ± 0.6 Ma) (7), the Nantuo Formation in South China (636.3 ± 4.9 Ma) (8), which underlies the cap carbonate of the basal Doushantuo Formation (635.2 ± 0.2 Ma) (9); and other glacial deposits around the globe, including the Ice Brook Formation in northwestern Canada (10). In contrast, a paucity of robust paleomagnetic poles and precise age constraints from volcanic rocks directly interbedded with early Cryogenian glacial deposits has precluded tests of the snowball Earth hypothesis for the Sturtian glaciation. The global nature of the Sturtian glaciation has been inferred from the ubiquitous occurrence of glacial deposits that are stratigraphically below Marinoan diamictite units (10) as well as banded iron formation within these deposits (1).

1Department of Earth and Planetary Sciences, Harvard University, Cambridge, MA 02138, USA. 2Department of Geosciences, Boise State University, Boise, ID 83725, USA. 3Geological Survey of Canada, Whitehorse, YT Y1A 1B5, Canada. 4Department of Earth and Planetary Sciences, Washington University, St. Louis, MO 63130, USA. 5Department of Geosciences, Princeton University, Princeton, NJ 08544, USA. 6555 East Simpson Street, Jackson, WY 83001, USA.

*To whom correspondence should be addressed. E-mail: fmacdon@fas.harvard.edu

Fig. 1. Composite chemo- and lithostratigraphy of Neoproterozoic strata in northwestern Canada, including the Upper and Lower Tindir Groups (13), the Fifteenmile and Lower Mount Harper Group (table S2), the Little Dal Group (3), the Coates Lake Group (table S2), the Twitya Formation (7), and the Shaler Supergroup (18). Shaded area represents the Bitter Springs isotopic stage (3).
east through the Ogilvie and Weekeepeemee Mountains of Yukon, to the Mackenzie Mountains of the Northwest Territories, and north to Victoria Island of Nunavut (fig. S1A). Exposures in the Coal Creek inlier of the central Ogilvie Mountains consist of mixed carbonate and siliciclastic rocks of the upper Fifteenmile and Lower Mount Harper Groups (LMHG), bimodal volcanic rocks of the Mount Harper volcanic complex (MHVC) (12), and glacial diamicite of the Upper Mount Harper Group (UMHG) (fig. 1). A glacial origin for the UMHG is inferred from bed-penetrating dropstones with impact margins and outsized clasts in fine, laminated beds, and by striated clasts in exposures in the Hart River inlier of the eastern Ogilvie Mountains (fig. S2). Evidence for grounded ice is provided by glacial push structures and soft-sedimentary deformation (fig. S2). The UMHG and the iron-rich unit 2 of the Upper Tindir Group in the western Ogilvie Mountains are correlative with the Sturtian-age Sayunei Formation of the Rapitan Group in the Mackenzie Mountains (13, 14). The broad distribution of massive diamicite and stratified glacial deposits with coarse-grained ice-rafter debris in the Rapitan Group of the Northwest Territories (14, 15) and its correlatives in Yukon and Alaska (fig. S3) suggests the proximity to a marine ice grounding line.

The MHVC previously was dated with multigrain U-Pb ID-TIMS analyses at 751 ± 26 Ma (16). We collected a quartz-phryic rhyolite of member D from the same site (fig. S1) that yielded a weighted mean 206Pb/238U zircon date of 717.43 ± 0.14 Ma, interpreted as the eruptive age of this unit (fig. S4). This ~33-million-year age revision is likely due to inherited cores in the previously dated multigrain zircon fractions, resulting in an artificially old age. Below the MHVC, a green, flinty, bedded tuff within allogenic dolostone beds near the top of unit PF1a of the Fifteenmile Group yielded a weighted mean 206Pb/238U zircon date of 811.51 ± 0.25 Ma, interpreted as the time of deposition (fig. S4). Above the MHVC, a green to pink brecciated tuff within glacial deposits of the UMHG yielded a weighted mean 206Pb/238U zircon date of 716.47 ± 0.24 Ma, interpreted as the deposition age (fig. S4).

In the Minto inlier on Victoria Island (fig. S1), zircon and baddeleyite from gabbroic sills and dikes from the Franklin LIP previously were dated at 723 ± 2 Ma and 718 ± 2 Ma (17). Our Franklin LIP sample is from a coarse-grained diabase sill, >20 m thick, intruding the middle of the Wynnati Formation (18), which yielded a weighted mean 206Pb/238U baddeleyite date of 716.33 ± 0.54 Ma (fig. S4). We interpret the apparent discrepancy between our result and previous ages (17) as an artifact of comparing upper intercept and 206Pb/206Pb dates with our 206Pb/238U dates in light of recently recognized systematic error in the U decay constant ratio (11, 19).

This geochronology reveals the >3000-km extent of the Franklin LIP (~716.5 Ma), from the Yukon-Alaska border to Ellesmere Island, where mafic dikes have been dated at 716 ± 1 Ma (20). Although no evidence for prevolcanic extension and rifting is present on Victoria Island (21), conspicuous normal faulting exists within the LMHG and the lower suite of the MHVC (12), temporally linking the Franklin LIP to extension on the northwestern Laurentian margin.

Several paleomagnetic studies on strongly magnetized mafic dikes, sills, and lavas have demonstrated that the Franklin LIP was emplaced when northwestern Laurentia was within 10° of the equator (6, 22, 23). The dated sill and the sediments that it intrudes on Victoria Island yield paleomagnetic data that are consistent with the previous low-latitude results (18). The age of the tuff interbedded with glacial deposits of the UMHC, 716.47 ± 0.24 Ma, is indistinguishable from the date of the Franklin LIP, 716.33 ± 0.54 Ma. Therefore, grounded ice was present on the northwestern margin of Laurentia at ~716.5 Ma, when it was situated at equatorial latitudes. Climate models have long predicted that if the ice line advanced equatorward of ~30° to 40°, an ice-albedo feedback would drive global glaciation (24, 25). Thus, we conclude that the Sturtian glaciation at ~716.5 Ma was global in nature.

It is uncertain whether the Sturtian glacial epoch consisted of one discrete glaciation that lasted tens of millions of years, or multiple glacial episodes including the low-latitude glaciation at ~716.5 Ma. Prior to this study, minimum and maximum age constraints on the Sturtian glaciation were provided by a sample from South China dated at 662.9 ± 4.3 Ma (26) and the Leger Granite in Oman dated at 726 ± 1 Ma (27), respectively. We suggest that our age from member D of the MHVC, 717.43 ± 0.14 Ma, provides a maximum age constraint on the low-latitude Sturtian glaciation not only because glacial deposits have not been identified below member D, but also because models suggest extremely rapid ice advance once ice is below 30° latitude (24, 25), such that glaciation of equatorial latitudes should be synchronous around the globe. Evidence for a pre-Sturtian glaciation at ~750 Ma (referred to as the Kaigas glaciation) was reported in southern Namibia (28), Zambia (29), and northwestern China (30). However, at these localities the contact relationship of the purported glacial deposit with the dated unit and the glacial origin of the deposit are sus-

![Fig. 2. Neoproterozoic composite carbonate δ13C chemostratigraphy with U-Pb ID-TIMS ages that are directly linked to isotopic profiles (12). Bars indicate the time spans of fossil assemblages representing eukaryotic crown groups. Asterisks indicate fossil groups of uncertain taxonomic affinity. Bars faded upward reflect uncertainty in the minimum age constraint; bars faded downward reflect uncertainty in the maximum age constraint. Dashes with question marks indicate that earlier records have not been proposed but the relationships between these fossils and the crown groups are uncertain.](image-url)
The Role of Sulfuric Acid in Atmospheric Nucleation

Mikko Sipilä,1,2,3,4 Torsten Berndt,1 Tuukka Petäjä,2 David Brus,3,5 Joonas Vanhanen,2 Frank Stratmann,1 Johanna Patokoski,2 Roy L. Mauldin III,4 Anti-Pekka Hyvärinen,5 Heikki Lihtavainen,5 Markku Kulmala2,6

Nucleation is a fundamental step in atmospheric new-particle formation. However, laboratory experiments on nucleation have systematically failed to demonstrate sulfuric acid particle formation rates as high as those necessary to account for ambient atmospheric concentrations, and the role of sulfuric acid in atmospheric nucleation has remained a mystery. Here, we report measurements of new particles (with diameters of approximately 1.5 nanometers) observed immediately after their formation at atmospherically relevant sulfuric acid concentrations. Furthermore, we show that correlations between measured nucleation rates and sulfuric acid concentrations suggest that freshly formed particles contain one to two sulfuric acid molecules, a number consistent with assumptions that are based on atmospheric observations. Incorporation of these findings into global models should improve the understanding of the impact of secondary particle formation on climate.

Nucleation of particles in the atmosphere has been observed to be strongly dependent on the abundance of sulfuric acid (H2SO4) (1–4). Sulfur dioxide (SO2), the precursor of H2SO4, has both natural and anthropogenic sources. Anthropogenic SO2 emissions can have large indirect effects on climate if H2SO4 is responsible for atmospheric nucleation, but laboratory experiments have systematically failed to reproduce ambient new-particle formation rates as well as the nucleation rate dependence on sulfuric acid.

References
11. See supporting material on Science Online.